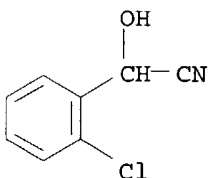


2 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2004 ACS on STN
 RN 13312-84-0 REGISTRY
 CN Benzeneacetonitrile, 2-chloro- α -hydroxy- (9CI) (CA INDEX NAME)
 OTHER CA INDEX NAMES:
 CN Mandelonitrile, o-chloro- (6CI, 7CI, 8CI)
 OTHER NAMES:
 CN (+)-2-Chloromandelonitrile
 CN (o-Chlorophenyl)glycolonitrile
 CN 2-Chlorobenzaldehyde cyanohydrin
 CN **2-Chloromandelonitrile**
 CN o-Chlorobenzaldehyde cyanohydrin
 CN o-Chloromandelonitrile
 FS 3D CONCORD
 DR 137766-65-5
 MF C8 H6 Cl N O
 LC STN Files: BEILSTEIN*, CA, CAOLD, CAPLUS, CASREACT, SPECINFO, TOXCENTER,
 USPATFULL
 (*File contains numerically searchable property data)
 DT.CA Caplus document type: Journal; Patent
 RL.P Roles from patents: ANST (Analytical study); BIOL (Biological study);
 PREP (Preparation); PROC (Process); RACT (Reactant or reagent); USES
 (Uses); NORL (No role in record)
 RL.NP Roles from non-patents: ANST (Analytical study); BIOL (Biological
 study); PREP (Preparation); PROC (Process); PRP (Properties); RACT
 (Reactant or reagent); NORL (No role in record)



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

36 REFERENCES IN FILE CA (1907 TO DATE)
 36 REFERENCES IN FILE CAPLUS (1907 TO DATE)
 3 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

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L3 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2004 ACS on STN
 RN 10421-85-9 REGISTRY
 CN Benzeneacetic acid, 2-chloro- α -hydroxy- (9CI) (CA INDEX NAME)
 OTHER CA INDEX NAMES:
 CN Mandelic acid, o-chloro- (6CI, 7CI, 8CI)
 OTHER NAMES:
 CN (+)-2-Hydroxy-2-(2-chlorophenyl)acetic acid
 CN (+)-o-Chloromandelic acid
 CN (2-Chlorophenyl)glycolic acid
 CN (2-Chlorophenyl)hydroxyacetic acid
 CN 2-Chloro- α -hydroxybenzeneacetic acid
 CN **2-Chloromandelic acid**
 CN NSC 31401
 CN o-Chloromandelic acid
 FS 3D CONCORD
 DR 52923-23-6
 MF C8 H7 Cl O3
 CI COM
 LC STN Files: BEILSTEIN*, BIOSIS, CA, CAOLD, CAPLUS, CASREACT, CHEMCATS,
 CHEMLIST, CSCHEM, IFICDB, IFIPAT, IFIUDB, IPA, SPECINFO, TOXCENTER,
 USPAT2, USPATFULL

(*File contains numerically searchable property data)

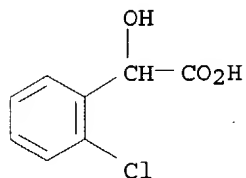
• Other Sources: EINECS**

(**Enter CHEMLIST File for up-to-date regulatory information)

DT.CA Caplus document type: Journal; Patent

RL.P Roles from patents: BIOL (Biological study); PREP (Preparation); PROC (Process); RACT (Reactant or reagent); NORL (No role in record)

RL.NP Roles from non-patents: ANST (Analytical study); BIOL (Biological study); FORM (Formation, nonpreparative); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses); NORL (No role in record)



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

66 REFERENCES IN FILE CA (1907 TO DATE)

66 REFERENCES IN FILE CAPLUS (1907 TO DATE)

5 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

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(FILE 'HOME' ENTERED AT 13:39:04 ON 29 SEP 2004)

FILE 'REGISTRY' ENTERED AT 13:39:15 ON 29 SEP 2004

1 0 S R-2-CHLOROMANDELONITRILE/CN
2 1 S 2-CHLOROMANDELONITRILE/CN
3 1 S 2-CHLOROMANDELIC ACID/CN
4 1 S 2-CHLOROMANDELIC ACID/CN

FILE 'CAPLUS' ENTERED AT 13:41:44 ON 29 SEP 2004

5 13 S 10421-85-9/PREP
6 6 S 10421-85-9/PROC
7 0 S 10421-85-9/PUR
8 19 S L5 OR L6
S L8 AND 13312-84-0/REG#

FILE 'REGISTRY' ENTERED AT 13:43:08 ON 29 SEP 2004

9 1 S 13312-84-0/RN

FILE 'CAPLUS' ENTERED AT 13:43:08 ON 29 SEP 2004

10 36 S L9
11 2 S L8 AND L10
12 1 S L11 AND OPTICAL PUR?

> s l10 and acid and hydroly?

3874803 ACID

569414 HYDROLY?

13 8 L10 AND ACID AND HYDROLY?

> s l10 and acid

3874803 ACID

14 23 L10 AND ACID

> s l10 and hydroly?

569414 HYDROLY?

15 9 L10 AND HYDROLY?

> s l15 and py<2000

19731428 PY<2000

16 6 L15 AND PY<2000

> d ibib abs hitstr

16 ANSWER 1 OF 6 CAPLUS COPYRIGHT 2004 ACS on STN

CESSION NUMBER: 1996:367768 CAPLUS

OCUMENT NUMBER: 125:32072

ITILE: Method of producing optically active α -hydroxy
acid or α -hydroxyamide

INVENTOR(S): Tamura, Koji

ATENT ASSIGNEE(S): Nitto Chemical Industry Co., Ltd., Japan

OURCE: Eur. Pat. Appl., 10 pp.

CODEN: EPXXDW

OCUMENT TYPE: Patent

ANGUAGE: English

AMILY ACC. NUM. COUNT: 1

ATENT INFORMATION:

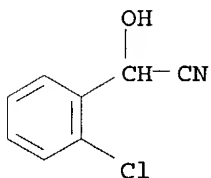
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 711836	A1	19960515	EP 1995-307976	19951108 <--
EP 711836	B1	20000202		
R: DE, FR, GB				
JP 08131188	A2	19960528	JP 1994-299109	19941109 <--
JP 3119468	B2	20001218		
US 5736385	A	19980407	US 1995-556085	19951109 <--
PRIORITY APPLN. INFO.:			JP 1994-299109	A 19941109

OTHER SOURCE(S): CASREACT 125:32072; MARPAT 125:32072

B A reaction system, wherein a cyanohydrin is converted to an optically
active α -hydroxy acid or α -hydroxyamide via a treatment in a
reaction tank with a microorganism, is provided with an automatic

cyanohydrin controller comprising a cyano ion detector, a regulator, and a cyanohydrin supplier linked thereto. The reaction is performed while automatically controlling the cyanohydrin concentration. Thus cyanohydrin can be supplied under automatic control at a relatively low and constant concentration on the basis of its consumption ratio. The reaction rate of the catalyst can be continuously regarded as the rate-limiting factor. As a result, a decrease in the enzymic activity during the reaction can be suppressed and an optically active α -hydroxy acid or α -hydroxyamide can be efficiently obtained at a high yield.

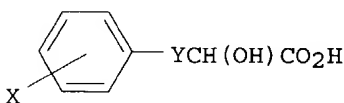
IT 13312-84-0, 2-Chloromandelonitrile
 RL: BPR (Biological process); BSU (Biological study, unclassified); RCT (Reactant); BIOL (Biological study); PROC (Process); RACT (Reactant or reagent)
 (microbial production of optically active α -hydroxy acids or α -hydroxyamides from α -hydroxy nitriles)
 RN 13312-84-0 CAPLUS
 CN Benzeneacetonitrile, 2-chloro- α -hydroxy- (9CI) (CA INDEX NAME)



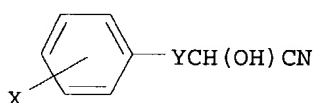
=> d ibib 2-6 abs hitstr

L16 ANSWER 2 OF 6 CAPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 1994:699313 CAPLUS
 DOCUMENT NUMBER: 121:299313
 TITLE: Process for producing optically active α -hydroxycarboxylic acid having phenyl group.
 INVENTOR(S): Hashimoto, Yoshihiro; Endo, Takakazu; Tamura, Koji; Hirata, Yuji
 PATENT ASSIGNEE(S): Nitto Chemical Industry Co., Ltd., Japan
 SOURCE: Eur. Pat. Appl., 10 pp.
 CODEN: EPXXDW
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 610048	A2	19940810	EP 1994-300704	19940131 <--
EP 610048	B1	19990922		
R: DE, FR, GB				
JP 06237789	A2	19940830	JP 1993-37275	19930203 <--
US 5580765	A	19961203	US 1994-191164	19940203 <--
PRIORITY APPLN. INFO.:			JP 1993-37275	19930203
OTHER SOURCE(S):	CASREACT 121:299313; MARPAT 121:299313			
GI				



I



II

AB A biol. process predominantly produces optically active α -hydroxycarboxylic acids I (X = H, OH, or C1-3 aliphatic saturated alkoxy, thioalkyl, halo, Ph, phenoxy, amino, or nitro; Y = (CH₂)_n where n = 0-2)

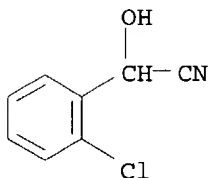
by asym. **hydrolysis** of a a racemic α -hydroxynitrile II (X and n as above) or a mixture of an aldehyde corresponding to the nitrile and prussic acid in a neutral to basic aqueous medium by the microorganism *Gordona terrae*, isolated from soil. A desired optically active α -hydroxycarboxylic acid having a Ph group can be obtained quant. at a high optical purity. Thus, a *G. terrae* suspension with OD630 = 20 in a pH = 8.2 phosphate buffer at 30C was shaken for 96 h with 10 mM phenylaldehyde + 10 mM KCN. The yield of 3-phenyllactic acid was 7.5 mM (75%) and the optical purity (L-form) was 63%.

IT 13312-84-0, 2-Chloromandelonitrile

RL: RCT (Reactant); RACT (Reactant or reagent)
(asym. **hydrolysis** of, by *Gordona terrae*)

RN 13312-84-0 CAPLUS

CN Benzeneacetonitrile, 2-chloro- α -hydroxy- (9CI) (CA INDEX NAME)



L16 ANSWER 3 OF 6 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 1992:5338 CAPLUS

DOCUMENT NUMBER: 116:5338

TITLE: Enzymic process for producing R(-)-mandelic acid and derivatives thereof

INVENTOR(S): Endo, Takakazu; Tamura, Koji

PATENT ASSIGNEE(S): Nitto Chemical Industry Co., Ltd., Japan

SOURCE: Eur. Pat. Appl., 28 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 449648	A2	19911002	EP 1991-302802	19910328 <--
EP 449648	A3	19920722		
EP 449648	B1	19990512		
R: DE, FR, GB				
JP 04099495	A2	19920331	JP 1990-214914	19900816 <--
JP 04099496	A2	19920331	JP 1990-214915	19900816 <--
JP 2698936	B2	19980119		
JP 04218385	A2	19920807	JP 1991-89189	19910329 <--
US 5223416	A	19930629	US 1991-677175	19910329 <--
PRIORITY APPLN. INFO.:				JP 1990-80694 19900330
				JP 1990-214914 19900816
				JP 1990-214915 19900816

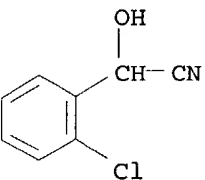
OTHER SOURCE(S): CASREACT 116:5338

AB A process for producing R(-)-mandelic acid or a derivative thereof comprises subjecting (i) R,S-mandelonitrile or a derivative thereof or (ii) a mixture of prussic acid and benzaldehyde or a derivative of benzaldehyde to the action of *Aureobacterium*, *Pseudomonas*, *Casheobacter*, *Alcaligenes*, *Acinetobacter*, *Brevibacterium*, *Nocardia*, and *Bacillus* or treated cells thereof, which microorganism is capable of stereospecifically **hydrolyzing** a nitrile group of the R,S-mandelonitrile or derivative, in a neutral or basic aqueous reaction system, to produce the R(-)-mandelic acid. Thus, racemic mandelonitrile was incubated with *Alcaligenes* BC24 for 16-24 h at 30° and pH 7.5. R(-) Mandelic acid was produced in 98% yield with an optical purity of 100%.

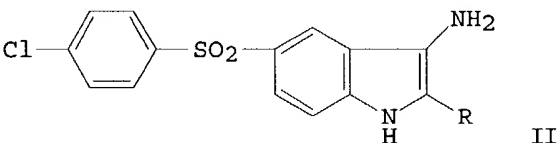
IT 13312-84-0

RL: RCT (Reactant); RACT (Reactant or reagent)
(asym. **hydrolysis** of, with microorganisms)

RN 13312-84-0 CAPLUS
CN Benzeneacetonitrile, 2-chloro- α -hydroxy- (9CI) (CA INDEX NAME)



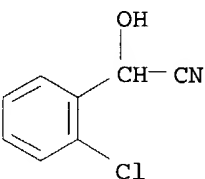
L16 ANSWER 4 OF 6 CAPLUS COPYRIGHT 2004 ACS on STN
ACCESSION NUMBER: 1988:112128 CAPLUS
DOCUMENT NUMBER: 108:112128
TITLE: Preparation and antimicrobial activity of
p-chloro-p'-(α -carbamoylbenzylamino)diphenyl
sulfones and 3-amino-2-aryl-5-p-
chlorophenylsulfonylindoles
AUTHOR(S): Mehta, K. J.; Prekh, H. H.; Parikh, A. R.
CORPORATE SOURCE: Dep. Chem., Saurashtra Univ., Rajkot, India
SOURCE: Acta Ciencia Indica, Chemistry (1985),
11(3), 187-90
CODEN: ACICDV; ISSN: 0253-7338
DOCUMENT TYPE: Journal
LANGUAGE: English
GI



AB 4-ClC6H4SO2C6H4NHCHRR1 (I, R = Ph, 2-HOC6H4, 3-HOC6H4, 4-HOC6H4, MeOC6H4, 2-ClC6H4, 4-ClC6H4, 3,4-MeO(HO)C6H3, PhCH:CH, 4-O2NC6H4, 2-furyl; R1 = cyano) were prepared by treating RCH(OH)CN with 4-ClC6H4SO2C6H4NH2-4. Treatment of I (R1 = cyano) with H2SO4 for 2 days gave I (R1 = CONH2) whereas after 7 days the indoles II were obtained. I (R1 = CONH2) have bactericidal activity against Staphylococcus aureus and Escherichia coli (no data).

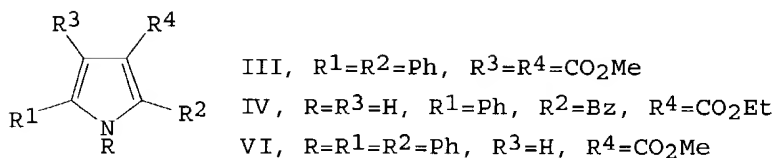
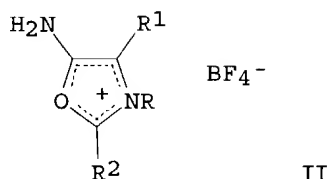
IT 13312-84-0
RL: RCT (Reactant); RACT (Reactant or reagent)
(reaction of, with aminochlorodiphenyl sulfone)

RN 13312-84-0 CAPLUS
CN Benzeneacetonitrile, 2-chloro- α -hydroxy- (9CI) (CA INDEX NAME)



L16 ANSWER 5 OF 6 CAPLUS COPYRIGHT 2004 ACS on STN
ACCESSION NUMBER: 1980:180928 CAPLUS
DOCUMENT NUMBER: 92:180928
TITLE: Synthetic uses of open-chain analogs of Reissert

compounds
 AUTHOR(S): McEwen, William E.; Grossi, Anthony V.; MacDonald, Russell J.; Stamegna, Andrew P.
 CORPORATE SOURCE: Dep. Chem., Univ. Massachusetts, Amherst, MA, 01003, USA
 SOURCE: Journal of Organic Chemistry (1980), 45(7), 1301-8
 CODEN: JOCEAH; ISSN: 0022-3263
 DOCUMENT TYPE: Journal
 LANGUAGE: English
 OTHER SOURCE(S): CASREACT 92:180928
 GI



AB Open-chain analogs, RN(COR₂)CHR₁CN (I, R = Ph, PhCH₂, p-ClC₆H₄, p-MeOC₆H₄, Me(CH₂)₅, cyclohexyl; R₁ = Ph, H, o-, m-, p-ClC₆H₄, 3,4-(MeO)₂C₆H₃, o-, m-MeOC₆H₄, Bu; R₂ = Ph, Me), of Reissert compds. are obtained by reaction of R₁CH(OH)CN with RNH₂, the resulting aminonitriles, RNHCHR₁CN, then being acylated. Hydrofluoroborate salts, II, of I, are prepared by reaction with fluoroboric acid in HOAc. The salts, II, undergo 1,3-dipolar addition reactions with reactive alkynes to give substituted pyrroles and with Et acrylate to give a different type of substituted pyrrole, the initial step in this instance being a Diels-Alder reaction. Thus, addition of MeO₂CC.tplbond.CCO₂Me to II (R₁ = R₂ = Ph) gave III (R = Ph, m-ClC₆H₄, p-MeOC₆H₄, PhCH₂); and addition of H₂C:CHCO₂Et to II (R = R₁ = R₂ = Ph) gave IV. I also undergo base-catalyzed reactions, such as alkylation with R₅Br to provide R₂CONRCR₁R₅CN (R₅ = PhCH₂, Bu, α-naphthylmethyl, R-R₂ = as above), which, in turn, undergo cleavage reactions in ethanolic alkali to give ketones R₁R₅CO. A conjugate addition reaction of the anion BzNPhC-PhCN (V) to Me acrylate to give, after subsequent steps, VI was demonstrated. α-Anilino ketones, PhNHCHRCOR₁, result when the anion V is treated with aldehydes, the initial reaction mixts. being subjected to subsequent alkaline **hydrolysis**. Finally, N-benzyl Reissert analogs give desoxybenzoins plus benzonitriles on treatment with NaH in THF.

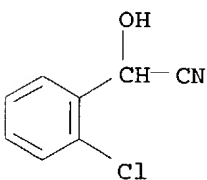
IT 13312-84-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reaction with amines, open-chain analogs of Reissert compds. from)

RN 13312-84-0 CAPLUS

CN Benzeneacetone nitrile, 2-chloro-α-hydroxy- (9CI) (CA INDEX NAME)



ACCESSION NUMBER: 1949:2594 CAPLUS
DOCUMENT NUMBER: 43:2594
ORIGINAL REFERENCE NO.: 43:605b-i,606a-c
TITLE: Preparation of the halophenylacetic acids
AUTHOR(S): Campbell, Neil; McKail, John E.
SOURCE: Journal of the Chemical Society, Abstracts (1948) 1251-5
CODEN: JCSAAZ; ISSN: 0590-9791
DOCUMENT TYPE: Journal
LANGUAGE: Unavailable
OTHER SOURCE(S): CASREACT 43:2594

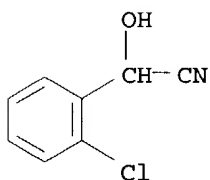
AB Granacher's synthesis (C.A. 16, 3898; 17, 2424) has been applied to the preparation of $\text{XC}_6\text{H}_4\text{CH}_2\text{CO}_2\text{H}$; the method depends on a satisfactory preparation of rhodanine (I), which is discussed. The yield of I depends upon the yield of $\text{H}_2\text{NCSSNH}_4$ (II) in the 1st stage of the reaction; the yield of II, as compared with that of $\text{CS}(\text{SNH}_4)_2$ (III), is a function of the NH_3 concentration which, in turn, is dependent on the temperature. The more concentrated the NH_3 solution, the greater is the yield of III; the best yield of II is obtained at 10-15°; the yield of I from II is 44%. When the temperature is kept below 0°, III is the main product. Methods of distinguishing between II and III are given. Mol. quantities of aldehyde and I in AcOH (5 cc. per g. aldehyde), refluxed 0.5 h. with fused AcONa (twice the weight of I), give the following benzylidenerhodanines (IV): o-Cl, pale yellow, m. 192°, 97% (Andreasch, C.A. 22, 3410, gives 169°); m-Cl, pale yellow, m. 233°, 93%; p-Cl, yellow, m. 231-2°, 93%; o-Br, orange, m. 183.5°, 80%; m-Br, yellow, m. 238°, 90%; p-Br, yellow, m. 237-8°, 84%. IV were transformed into β -phenyl- α -thiopyruvic acids (V) by heating with 8 cc. 8% NaOH (per g. IV) at 50-5° until a clear or nearly clear solution resulted, cooling in an ice-salt bath, and acidifying rapidly with 3 N HCl ; the crude acid in cold EtOH is precipitated with 1-2 vols. cold H_2O and recrystd. from MeOH , petr. ether, etc.; however, the crude acids were used in the next step. o-Cl, m. 134-5°, 72% (A. gives 119-20°); m-Cl, straw-colored, m. 134°, 84%; p-Cl, yellow, m. 169-71°, 84% [a byproduct is probably α, α' -dithiobis(m-chlorocinnamic acid), yellow, m. 221-2°]; o-Br, lemon-yellow, m. 142-3°, 70%; m-Br, pale yellow, m. 133-4°, 81%; p-Br, m. 165-80°, 75% (could not be purified further). α -Isonitroso- β -(halophenyl)propionic acids (VI) were prepared from V by refluxing (about 0.5 h.) in alc. containing 3 mols. NH_2OH (until H_2S evolution ceases); the crude acid is precipitated from dilute NaOH solution with concentrated HCl . o-Cl, m. 156°, 83%; m-Cl, m. 149° (decomposition), 100%; p-Cl, m. 170° (decomposition) (1 sample m. 182°), 100%; o-Br, m. 150° (decomposition), 100%; m-Br, m. 151°, 93%; p-Br, m. 173°, 85% (yields are of crude products). VI, refluxed 10 min. with Ac_2O (4 cc. per g. VI), the Ac_2O removed in vacuo, and the residue extracted with ether, give the (halophenyl)acetonitriles (VII): o-Cl, b11 123-5°, 64%; m-Cl, b10 134-6°, 55%; p-Cl, b12 137-9°, m. 31-2°, 80%; o-Br, b13, 140-1°, 88%; m-Br, b10 145-7°, m. 27-8°, 70%; p-Br, b10-12 152-6°, m. 48°, 72%. The over-all yields of the VII from the aldehydes were: o-, m-, and p-Cl, 57, 47, 62%; o-, m-, and p-Br, 49, 44, 38%; further losses, sometimes considerable, occur in the next step. VII were hydrolyzed by boiling with 60% H_2SO_4 or, preferably, with 20% EtOH-KOH , giving $\text{RC}_6\text{H}_4\text{CH}_2\text{CO}_2\text{H}$ (R given): o-Cl, m. 93-5° (p-nitrobenzyl ester, m. 70-1°); m-Cl, m. 77° (p-nitrobenzyl ester, m. 74-5°; p-toluidide, m. 138°); p-Cl, m. 104-6° (p-nitrobenzyl ester, m. 117°); o-Br, m. 104-5° (p-nitrobenzyl ester, m. 74-5°; p-toluidide, m. 183-4°; anilide, m. 153-4°); m-Br, m. 102-3° (p-nitrobenzyl ester, m. 75-6°; p-toluidide, m. 135°); p-Br, m. 113-15° (p-nitrobenzyl ester, m. 128-9°; anilide, m. 174-6°; p-toluidide, m. 203°). The results show that this method leaves much to be desired. The crystalline compound from 20 g. p- $\text{BrC}_6\text{H}_4\text{CHO}$, 80 cc. NaHSO_3 , and 5 cc. EtOH , stirred 2 h. with 10 g. KCN in 20 cc. H_2O , gives 12.5 g. p-bromomandelonitrile, m. 78-9°; 11 g. and 46 cc. HI (d. 1.94), refluxed 1 h. give 1.5 g.

^ p-BrC₆H₄CH₂CO₂H; an unknown compound, m. 126-7°, is a byproduct.
 • o-ClC₆H₄CHO, through o-ClC₆H₄CH(OH)CN, yields the benzoate
 (no properties given); refluxing with Pt black in tetralin did not give
 o-ClC₆H₄CH₂CN. p-MeOC₆H₄CH₂CN was obtained in 39% yield (yield of
 intermediate benzoate 74 and 92% in 2 expts.). Other methods were tried
 without much success. A mixture of o- and p-BrC₆H₄CH₂CO₂H could not
 be separated by fractional distillation of the acid chlorides or Et esters;
 chromatog. separation of the anilides, p-toluidides, and 2-naphthalides was
 only partially successful (they fluoresce in C₆H₆ but not on the Al₂O₃
 column). α-(o-Bromophenyl)aceto-2-naphthalide, m.
 188-9°; p-isomer, m. 203-4°. α-Phenylaceto-2-
 naphthalide m. 162-3°. In the preparation of the naphthalides by
 heating the acids with 2-C₁₀H₇NH₂, (2-C₁₀H₇)₂NH is obtained, the catalyst
 presumably being the halo acid. 2-C₁₀H₇NH₂ and PhNH₂ give 2-C₁₀H₇NHPh,
 and p-MeC₆H₄NH₂ gives 2-C₁₀H₇NHC₆H₄Me-p, but o- or m-ClC₆H₄NH₂
 gives only (2-C₁₀H₇)₂NH. 2-C₁₀H₇NH₂ and its derivs. are more strongly
 adsorbed than the corresponding 1-derivs.

IT 13312-84-0, Mandelonitrile, o-chloro-
 (preparation of)

RN 13312-84-0 CAPLUS

CN Benzeneacetonitrile, 2-chloro-α-hydroxy- (9CI) (CA INDEX NAME)



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Refine Search

Search Results -

Terms	Documents
L11 and cyanohydrin and hydroly\$6	3

Database:

US Pre-Grant Publication Full-Text Database
 US Patents Full-Text Database
 US OCR Full-Text Database
 EPO Abstracts Database
 JPO Abstracts Database
 Derwent World Patents Index
 IBM Technical Disclosure Bulletins

Search:

L14

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DATE: Wednesday, September 29, 2004 [Printable Copy](#) [Create Case](#)

<u>Set Name</u> side by side	<u>Query</u>	<u>Hit Count</u>	<u>Set Name</u> result set
<i>DB=PGPB,USPT,USOC,EPAB,JPAB,DWPI,TDBD; PLUR=YES; OP=ADJ</i>			
<u>L14</u>	l11 and cyanohydrin and hydroly\$6	3	<u>L14</u>
<u>L13</u>	l11 and cyanohydrin	3	<u>L13</u>
<u>L12</u>	L11 and 562/\$	11	<u>L12</u>
<u>L11</u>	L10 and rate	42	<u>L11</u>
<u>L10</u>	L9 and cool\$5 and crystal\$5	68	<u>L10</u>
<u>L9</u>	L8 and optically pure	117	<u>L9</u>
<u>L8</u>	hydroxycarboxylic acid	15476	<u>L8</u>
<u>L7</u>	L1 and optically pure	105	<u>L7</u>
<i>DB=PGPB,USPT; PLUR=YES; OP=ADJ</i>			
<u>L6</u>	L1 and optically pure	105	<u>L6</u>
<u>L5</u>	L4 and rate	0	<u>L5</u>
<u>L4</u>	L3 and cool\$5 and crystal\$5	2	<u>L4</u>
<u>L3</u>	L2 and optically pure	4	<u>L3</u>
<u>L2</u>	hydroxycarboxylic acid.ti.	139	<u>L2</u>

L1 hydroxycarboxylic acid

10088 L1

END OF SEARCH HISTORY

Hit List

Clear

Generate Collection

Print

Fwd Refs

Bkwd Refs

Generate OACS

Search Results - Record(s) 1 through 3 of 3 returned.

☐ 1. Document ID: US 20030215859 A1

Using default format because multiple data bases are involved.

L14: Entry 1 of 3

File: PGPB

Nov 20, 2003

PGPUB-DOCUMENT-NUMBER: 20030215859

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20030215859 A1

TITLE: DNA shuffling of monooxygenase genes for production of industrial chemicals

PUBLICATION-DATE: November 20, 2003

INVENTOR-INFORMATION:

NAME	CITY	STATE	COUNTRY	RULE-47
Affholter, Joseph A.	Zephyr Cove	NV	US	
Davis, S. Christopher	San Francisco	CA	US	
Selifonov, Sergey A.	Plymouth	MN	US	

US-CL-CURRENT: 435/6; 435/189, 435/320.1, 435/325, 435/7.1

Full	Title	Citation	Front	Review	Classification	Date	Reference	Sequences	Attachments	Claims	KMOC	Draw. De
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☐ 2. Document ID: US 6605430 B1

L14: Entry 2 of 3

File: USPT

Aug 12, 2003

US-PAT-NO: 6605430

DOCUMENT-IDENTIFIER: US 6605430 B1

TITLE: DNA shuffling of monooxygenase genes for production of industrial chemicals

Full	Title	Citation	Front	Review	Classification	Date	Reference	Sequences	Attachments	Claims	KMOC	Draw. De
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☐ 3. Document ID: US 4517194 A

L14: Entry 3 of 3

File: USPT

May 14, 1985

US-PAT-NO: 4517194

DOCUMENT-IDENTIFIER: US 4517194 A

TITLE: Azolylmandelic acid derivatives and use thereof for controlling

microorganisms

Full	Title	Citation	Front	Review	Classification	Date	Reference	Sequences	Attachments	Claims	KWIC	Draw. Data
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Clear	Generate Collection	Print	Fwd Refs	Bkwd Refs	Generate OACS
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Terms	Documents
L11 and cyanohydrin and hydroly\$6	3

Display Format: [Previous Page](#)[Next Page](#)[Go to Doc#](#)

Hit List

Clear	Generate Collection	Print	Fwd Refs	Bkwd Refs
Generate OACS				

Search Results - Record(s) 1 through 10 of 11 returned.

☐ 1. Document ID: US 20010051747 A1

Using default format because multiple data bases are involved.

L12: Entry 1 of 11

File: PGPB

Dec 13, 2001

PGPUB-DOCUMENT-NUMBER: 20010051747

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20010051747 A1

TITLE: Process for the separation of a mixture of enantiomers

PUBLICATION-DATE: December 13, 2001

INVENTOR-INFORMATION:

NAME	CITY	STATE	COUNTRY	RULE-47
Vries, Ton R.	Groningen		NL	
Wijnberg, Hans	Haren		NL	
Echten, Erik Van	Assen		NL	
Hulshof, Lumbertus A.	Baarlo		NL	
Broxterman, Quirinus B.	Sittard		NL	

US-CL-CURRENT: 562/401

Full	Title	Citation	Front	Review	Classification	Date	Reference	Sequences	Attachments	Claims	KWIC	Draw. De
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☐ 2. Document ID: US 6235927 B1

L12: Entry 2 of 11

File: USPT

May 22, 2001

US-PAT-NO: 6235927

DOCUMENT-IDENTIFIER: US 6235927 B1

**** See image for Certificate of Correction ****

TITLE: Process for the separation of a mixture of enantiomers

Full	Title	Citation	Front	Review	Classification	Date	Reference	Sequences	Attachments	Claims	KWIC	Draw. De
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☐ 3. Document ID: US 5986074 A

L12: Entry 3 of 11

File: USPT

Nov 16, 1999

US-PAT-NO: 5986074

DOCUMENT-IDENTIFIER: US 5986074 A

**** See image for Certificate of Correction ****

TITLE: Metal chelates as pharmaceutical imaging agents, processes of making such and uses thereof

Full	Title	Citation	Front	Review	Classification	Date	Reference	Sequence	Attachment	Claims	KWIC	Draw. De
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☐ 4. Document ID: US 5955053 A

L12: Entry 4 of 11

File: USPT

Sep 21, 1999

US-PAT-NO: 5955053

DOCUMENT-IDENTIFIER: US 5955053 A

**** See image for Certificate of Correction ****

TITLE: Metal chelates as pharmaceutical imaging agents, processes of making such and uses thereof

Full	Title	Citation	Front	Review	Classification	Date	Reference	Sequence	Attachment	Claims	KWIC	Draw. De
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☐ 5. Document ID: US 5350761 A

L12: Entry 5 of 11

File: USPT

Sep 27, 1994

US-PAT-NO: 5350761

DOCUMENT-IDENTIFIER: US 5350761 A

TITLE: Indolyl substituted hydroxylamine derivatives

Full	Title	Citation	Front	Review	Classification	Date	Reference	Sequence	Attachment	Claims	KWIC	Draw. De
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☐ 6. Document ID: US 5334600 A

L12: Entry 6 of 11

File: USPT

Aug 2, 1994

US-PAT-NO: 5334600

DOCUMENT-IDENTIFIER: US 5334600 A

TITLE: Isoquinolyl substituted hydroxylamine derivatives

Full	Title	Citation	Front	Review	Classification	Date	Reference	Sequence	Attachment	Claims	KWIC	Draw. De
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☐ 7. Document ID: US 5260316 A

L12: Entry 7 of 11

File: USPT

Nov 9, 1993

US-PAT-NO: 5260316

DOCUMENT-IDENTIFIER: US 5260316 A

TITLE: Isoquinolyl substituted hydroxylamine derivatives

Full	Title	Citation	Front	Review	Classification	Date	Reference	Sequences	Attachments	Claims	KWIC	Draw De
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☐ 8. Document ID: US 5246965 A

L12: Entry 8 of 11

File: USPT

Sep 21, 1993

US-PAT-NO: 5246965

DOCUMENT-IDENTIFIER: US 5246965 A

TITLE: Arylethers, their manufacture and methods of treatment

Full	Title	Citation	Front	Review	Classification	Date	Reference	Sequences	Attachments	Claims	KWIC	Draw De
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☐ 9. Document ID: US 4384879 A

L12: Entry 9 of 11

File: USPT

May 24, 1983

US-PAT-NO: 4384879

DOCUMENT-IDENTIFIER: US 4384879 A

TITLE: 4-(1H-Azolylmethyl)-1,3-dioxolan-5-one derivatives, production thereof and use thereof as growth regulators and/or microbicides

Full	Title	Citation	Front	Review	Classification	Date	Reference	Sequences	Attachments	Claims	KWIC	Draw De
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☐ 10. Document ID: US 3711528 A

L12: Entry 10 of 11

File: USPT

Jan 16, 1973

US-PAT-NO: 3711528

DOCUMENT-IDENTIFIER: US 3711528 A

TITLE: RACEMIC DIHYDRO-PGE AND RELATED COMPOUNDS

Full	Title	Citation	Front	Review	Classification	Date	Reference	Sequences	Attachments	Claims	KWIC	Draw De
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Terms	Documents
L11 and 562/\$	11

Display Format: